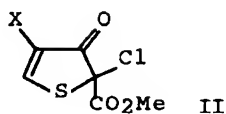
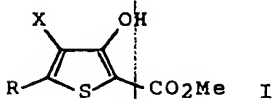


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ACCESSION NUMBER: 1988:131570 HCAPLUS COPYRIGHT 2006 ACS on STN
 Full-text
 DOCUMENT NUMBER: 108:131570
 TITLE: Process for the preparation of 3-hydroxy-5-(1-polyazolyl)-2-methoxycarbonylthiophenes, useful as pharmaceuticals or their intermediates
 INVENTOR(S): Corral Saleta, Carlos; Lissavetzky Diez, Jaime
 PATENT ASSIGNEE(S): Consejo Superior de Investigaciones Cientificas, Spain
 SOURCE: Span., 7 pp.
 CODEN: SPXXAD
 DOCUMENT TYPE: Patent
 LANGUAGE: Spanish
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
ES 547442	A1	19861116	ES 1985-547442	19850930
PRIORITY APPLN. INFO.:			ES 1985-547442	19850930



AB The title compds. (I; X = H, halo; R = heterocyclic polyazolo or benzopolyazolo), useful as intermediate or final products in the synthesis of new therapeutic agents (no data), are prepared by condensation with chloro(methoxycarbonyl)oxodihydrothiophenes II with 2 equivalent polyazole or benzopolyazole RH at room temperature. A solution of 0.009 mol II (X = H) in HOAc was treated with 0.018 mol pyrazole and kept at room temperature for 2 days to give crystalline I (X = H, R = 1-pyrazolyl).